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- (54) PLASTIC ARTICLES COMPRISING BIODEGRADABLE PHA COPOLYMERS

POLYHYDROXYALKANOAT-COPOLYMERE ENTHALTENDE ABBAUBARE KUNSTSTOFFARTIKEL

ARTICLES EN PLASTIQUE RENFERMANT DES COPOLYMERES PHA BIODEGRADABLES

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Description

TECHNICAL FIELD

5 [0001] The present invention relates to plastic articles other than a film comprising biodegradable PHA copolymers.

BACKGROUND

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[0002] Polymers find uses in a variety of plastic articles including films, sheets, fibers, foams, molded articles, adhesives and many other specialty products. For applications in the areas of packaging, agriculture, household goods and personal care products, prolymers usually have a short (less than 12 months) use cycle. For example, in food packaging, polymers play the role of a protective agent and are quickly disposed of after the contents are consmuned. Household products such as detergent bottles and diapsers are immediately discarded once the product is used.

[0003] The majority of this plastic material ends up in the solid waste stream, headed for rapidly vanishing and increasingly expensive landfill space. While some efforts at recycling have been made, the nature of polymers and the way they are produced and convented to products limits the number of possible recycling applications. Repeated processing of even pure polymers results in degradation of material and consequently poor mechanical properties. Different grades of chemically similar plastics (e.g., polyethylenes of different molecular weights, as used in milk jugs and grocery sacks) mixed upon collection can cause processing problems that make the reclaimed material inferior or nursable.

(1004) Absorbent article applications such as dispers, sanitary napkins, pantiliners and the like, involve several different types of plastics. In these cases, recycling is particularly costly because of the difficulty in separating the different types of plastics. In these cases, recycling is particularly costly because of the difficulty in separating the different components. Disposable products of this type generally comprise some sort of fluid-permeable besident materials, and absorbent core, and a fluid-impermeable backsheet materials. Heretofore, such absorbent structures have been prepared using, for example, topsheet materials prepared from woven, nor-woven, or prorous formed-film polyethene or polyproprise materials. Backsheet materials typically comprise flexible polyethylene sheets. Absorbent core materials typically comprise wood pulp filters or wood pulp filters in combination with absorbent gelling materials. Although such products largely comprise materials which would be expected utilimately to degrade, and although products of this type contribute only a very small percentage of the total solid waste materials generated by consumers each year, nevertheless, there is currently a proreview feed to devise such disposable products from materials which are composable.

30 [0005] A conventional disposable absorbent product is already to a large extent compostable. A typical disposable diager, for example, consists of about 80% of compostable materials, e.g., wood putp fibers, and the fike. In the composting process solied disposable absorbent articles are stredded and commingled with organic waste prior to the composting per se. After composting is complete, the non-compostable particles are screened out. In this manner even today's absorbent articles can successfully be processed in commercial composting plants.

[0005] Nevertheless, there is a need for reducing the amount of non-compostable materials in disposable absorbent articles. There is a particular need to replace polyethylene backsheets in absorbent articles with liquid impervious films of compostable material, because the backsheet is typically one of the largest non-compostable components of a conrentional disposable absorbent article.

[0007] In addition to being compostable, the films employed as backsheets for absorbent articles must satisfy many to other performance requirements. For example, the resists should be termoplastic such that conventional film processing methods can be employed. These methods include cast film and blown film extrusion of single leyer structures and cast or blown film coextrusion of multilayer structures. Other methods include extrusion coating of one material on one or both sides of a compostable substrate such as another film, a non-woren fabric, or a paper well.

[0008] Still other properties are essential in product converting operations where the films are used to fabricate absorbent articles. Properties such as tensile strength, tensile modulus, tear strength, and thermal softening point determine to all large extent how well all film will run on converting lines.

[0009] In addition to the aforementioned properties, still other properties are needed to meet the end user requirements of the absorbent article. Film properties such as impact strength, puncture strength and moisture transmission are important since hely influence the absorbent article is durability and containment while being worn.

[0010] Once the absorbent article is disposed of and enters a composting process, other properties become important. Regardless of whether incoming waste is preshredded or not, it is important that the film or large film fragments undergo an initial breakup to much smaller particles during the initial stages of composting. Otherwise, the films or large fragments may be screened out of the compost stream and may never become part of the final compost.

[0011] In the past, the biodegradability and physical properties of a variety of polyhydroxyalkanoates (PHAs) have been studied. Polyhydroxyalkanoates are polyeater compounds produced by a variety of microorgalmss, such as bacteria and algoe. While polyhydroxyalkanoates have been of general interest because of their biodegradable nature, their actual uses as a plastic material has been hampered by their thermal instability. For example, poly-3-hydroxyoutyrate (PHB) is a natural energy-schorage product of bacteria and algae, and is present in discrete granules within the cell

cytoplasm. However, unlike other biologically synthesized polymers such as proteins and polysaccharides, PHB is thermoplastic having a high degrace of crystalfinity and a well-defined melt temperature of about 180°C. Unfortunately, PHB becomes unstable and degrades at elevated temperatures near its melt temperature. Due to this thermal instability, commercial applications of PHB have been extremely limited.

[012] As a result, investigators have studied other polyhydroxyalkaneates such as poly(3-hydroxybulyrate-co-3-hydr

[0013] Due to the slow crystallization rate, a film made from PHBV will actic to itself even after cooling; a substantial fraction of the PHBV remains amorphous and tacky for long periods of time. In cast film operations, where the film is immediately cooled on chill rolls after leaving the film die, motten PHBV often sticks to the rolls restricting the speed at which the film can be processed, or even preventing the film from being collected. In blown films, residual tack of the PHBV causes the tubular film to sick to lister flater it has been cooled and collapsed or winding.

(014) U.S. Patent 4,880,932, Marthi et al., issued November 14, 1999, discloses a means of achieving a PHBV monolayer film for disper backsheet applications by coextruding the PHBV between two layers of sacrificial polymer, for example a polyolefin, stretching and orienting the multilayer film, and then stripping away the polyolefin layers after the PHBV has had time to crystalfize. The remaining PHBV film is then laminated to either water soluble films or water insoluble films outher polyolefins. Untofurnately, such drastic and cumbersome processing measures are necessary in an attempt to avoid the inherent difficulties associated with processing PHBV into films. [0015] Based on the foregoing, there is a need for plastic articles that can biodegradable in effect such biodegradable articles would facilitate the "recycling" of plastic articles into another usable product, topsoil, through composting. To salisfy this need, there is a preliminary need for a biodegradable polymer which is capable of being easily processed into a plastic article for use in a disposable product.

Objects of the Invention

[0016] It is an object of the present invention to provide plastic articles other than a film comprising a biodegradable polyhydroxyalkanoate (PHA).

SUMMARY

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[0017] The present invention relates to plastic articles other than a film comprising a biodegradable copolymer, wherein the copolymer comprises at least two randomly repeating monomer units wherein the first monomer unit has the structure

wherein R1 is H, or C1 or C2 alkyl, and n is 1 or 2; the second monomer unit has the structure

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and wherein at least 50% of the random repeating monomer units have the structure of the first monomer unit. Such plastic articles include films, sheets, libers, loams, molded articles, nomoven fabrics, elastomers, and adhesives, [0018] The present invention further relates to an absorbert article comprising a liquid previous topsheet, a biodegradable liquid impervious backsheet comprising a film comprising a biodegradable PHA, and an absorbert core positioned between the topsheet and the backsheet.

DETAILED DESCRIPTION

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[0019] The present invention answers the need for a biodegradable copolymer which is capable of being easily processed into a plastic article. The present invention further answers the need for disposable plastic articles with increased biodegradability and/or compostability.

[0020] As used herein, "ASTM" means American Society for Testing and Materials.

[0021] As used herein, "comprising" means that other steps and other ingredients which do not affect the end result can be added. This term encompasses the terms "consisting of" and "consisting essentially of".

[0022] As used herein, "alkyr" means a saturated carbon-containing chain which may be straight or branched; and substituted (mono- or poly-) or unsubstituted.

[0023] As used herein, 'alkenyl' means a carbon-containing chain which may be monounsaturated (i.e., one double bond in the chain) or polyunsaturated (i.e., two or more double bonds in the chain); straight or transhed; and substituted (mono- or poly) or unsubstituted.

[0024] As used herein, "PHA" means a polyhydroxyalkanoate of the present invention.

[0025] As used herein, "PHB" means the homopolymer poly-(3-hydroxybutyrate).

[0026] As used herein, "PHBV" means the copolymer poly(3-hydroxybutyrate-co-3-hydroxyvalerate).

[0027] As used herein, "PHBMV" means the copolymer poly(3-hydroxybutyrate-co-3-hydroxy-4-methylvalerate).

[0028] As used herein, "biodegradable" means the ability of a compound to ultimately be degraded completely into CO, and water or biomass by microorganisms and/or natural environmental factors.

[0029] As used herein, "compostable" means a material that meets the following three requirements: (1) the material is capable of being processed in a composting facility for solid waste; (2) if so processed, the material will end up in the final compost, and (3) if the compost is used in the soil, the meterial will ultimately biodegrade in the soil.

[0030] For example, a polymer film material present in solid waste submitted to a composting facility for processing does not necessarily end up in the final compost. Certain composting facilities subject the solid waste stream to air classification prior to further processing, in order to separate paper and other materials. A polymer film would most probably be separated from the solid waste stream in such an air classification and therefore not be processed in the compositing facility. Nevertheless, it may still be a "Compostable" material according to the above definition because it is "capable" of being processed in a composting facility.

[0031] The requirement that the material ends up in the final compost typically means that it undergoes a form of degradation in the composting process. Typically, the solid waste stream will be subjected to a shreading step in an early phase of the composting process. As a result, the polymer lim will be present as shreds rather than a sheet. In the final phase of the composting process, the finished compost will be subjected to a screening stor. Typically, the polymer streds will not pass through the screens if they have retained the size they had immediately after the shredding stop. The compostable materials of the present invention will have lost enough of their integrity during the composting process to allow partially degraded shreds to pass through the screens. However, it is conclevable that a composting actility might subject the solid waste stream to a very rigorous streading and a rather coarse screening, in which case nondegradable polymers like polyethylene would meet requirement (2). Therefore, meeting requirement (2) is not enough for a material to be compostable within the present definition.

[0032] What distinguishes the compostable material as defined herein from material like polyethylene is requirement.

(3) that the material ultimately biodegrade in the soil. This biodegradability requirement is not essential to the composting process or the use of composting soil. Solid waste and the compost resulting therefrom may contain all kinds of nonbiodegradable materials, for example, sand. However, to avoid a build up of man-made materials in the soil, it is required herein that such materials be fully biodegradable. By the same token, it is not at all necessary that this biodegradable

be fast. As long as the material itself and intermediate decomposition products are not toxic or otherwise harmful to the soil or crops, it is fully acceptable that their biodegradation takes several months or even years, since this requirement is present only to avoid an accumulation of man-made materials in the soil.

[0033] All copolymer composition ratios recited herein refer to mole ratios, unless specifically indicated otherwise.

[0034] The present invention relates to biodegradable copolymers which are surprisingly easy to process into plastic articles, as compared to the homopolymer PHB and copolymer PHBV.

[0035] As used herein, "plastic article" means a copolymer processed into a sheet, fiber, foam, molded article, nonwoven fabric, elastomer or adhesive.

[0036] PHAs useful for processing into plastic articles of the present invention comprise at least two randomly repeating monomer units (RRMU). The first RRMU has the structure

$$\begin{array}{c|c} & R^1 & O \\ \hline & O - CH - (CH_2)_n - C \end{array}$$

20 wherein R1 is H, or C1 or C2 alkyl, and n is 1 or 2. The second RRMU has the structure

[0037] In one embodiment of the present invention, at least about 50%, but less than 100%, of the RRMUs have the structure of the first RRMU; more preferably at least about 60%; more preferably at least about 70%; more preferably at least about 50%; more preferably at least about 50%; more preferably sit least about 50%.

[0038] When a PHA of the present invention is processed into a film, sheet, or soft elastic fiber, preferably from about 55% to about 95.9% of the RRM/Us have the structure of the first RRM/U unit, more preferably from about 75% to about 99%; more preferably 56% to about 95% about 95%.

[0033] When a PHA of the present invention is processed into a normal fiber or molded article (e.g., injected or blown molded) preferably from about 80% to about 99.5% of the first RRMUs have the structure of the first RRMU; more preferably from about 90% to most 99.5% more preferably from about 90% to about 99.5%.

40 [0040] When a PHA of the present invention is processed into an elastomer or an adhesive, preferably from about 50% to 85% of the RRMUs have the structure of the first RRMU.

[0041] When a PHA of the present invention is processed into a nonwoven, preferably from about 65% to about 99.5% of the RRMUs have the structure of the first RRMU; more preferably from about 90% to about 99.5%; more preferably still from about 95% to about 99.5%; more preferably still from about 95% to about 99.5%;

45 [0042] In one embodiment of the present invention, R1 is a C₁ alkyl and n is 1, thereby forming the monomeric repeat unit 3-hydroxybutyrate.

[0043] In another embodiment of the present invention, R¹ is a C₂ alkyl and n is 1, thereby forming the monomeric repeat unit 3-hydroxyvalerate.

[0044] In another embodiment of the present invention, R1 is H and n is 2, thereby forming the monomeric repeat unit 4-hydroxybutyrate.

[0045] In another embodiment of the present invention, R¹ is H and n is 1, thereby forming the monomeric repeat unit 3-hydroxypropionale.

[0046] In another embodiment, the copolymer useful in the present invention comprises one or more additional RRMUs having the structure

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$$\begin{array}{c|c} & & & & \\ \hline & & & & \\ \hline & O - CH - (CH_2)_m - C \end{array}$$

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wherein R^3 is H, or a C_1 , C_2 , C_3 , C_4 , C_6 , C_6 , C_7 , C_8 , C_9 , C_{10} , C_{11} , C_{12} , C_{12} , C_{12} , C_{13} , C_{15} , C_{15} , C_{15} , C_{15} or alkyriy, and m is 1 or 2; and wherein the additional RNNUL5 are not the same as the first RNNUL5 in the second RNNU. Preferably the copolymer comprises 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20 or more different RNNUs. 10047]. In a preferred embodiment of the present invention, R^3 is a C_1 , C_2 , C_3 , C_4 , C_6 , C_6 , C_7 , C_6 , C_9 ,

C₁₃ C₁₄ C₁₅ C₁₆ C₁₇ C₁₈ O C₁₉ any to an any an arrival and m is 1, thereby forming the monomeric [0048]. In a preferred embodiment of the present invention, R³ is a C₁ alkyl and m is 1, thereby forming the monomeric repeat unit 3-hydroxybulyrate.

5 [0049] In another embodiment of the present invention, R3 is a C₂ alkyl and m is 1, thereby forming the monomeric repeat unit 3-hydroxyvalerate.

[0050] In another embodiment of the present invention, R³ is H and m is 2, thereby forming the monomeric repeat unit 4-hydroxybutyrate.

[0051] In another embodiment of the present invention, R³ is H and m is 1, thereby forming the monomenic repeat or unit 3-hydroxypropionate.

[0052] Preferably, novel biodegradable PHAs of the present invention comprising two RRMUs have a first RRMU having the structure

wherein R1 is H, or C1 or C2 alkyl, and n is 1 or 2; and a second RRMU having the structure

wherein at least 50% of the RRMUs have the structure of the first RRMU. [0053] Preferably, novel biodegradable PHAs of the present invention comprising three RRMUs, have a first RRMU having the structure

$$\begin{array}{c|c} & R^1 & O \\ \hline -O-CH-(CH_2)_n-C \end{array}$$

wherein R1 is H, or C1 or C2 alkyl or alkenyl, and n is 1 or 2; a second RRMU having the structure

10 and a third RRMU having the structure

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wherein R³ is H, or a C₁, C₂, C₃, C₄, C₅, C₅, C₇, C₈, C₉, C₁₀, C₁₀, C₁₇, C₁₂, C₁₃, C₁₄, C₁₅, C₁₆, C₁₇, C₁₈, or C₁₉ alkyl or alkeryl; and m is 1 or 2, wherein at least 50% of the RRMUs have the structure of the first RRMU, and wherein the third RRMUs is not the same as the first randomly repeating monomer unit.

Synthesis of Biodegradable PHAs

[0054] The biodegradable PHAs of the present invention can be synthesized by synthetic chemical or biological based methods. A chemical approach involves the ring-opening polymerization of β-lactone monomers as described below. The catalysts or initiators used can be a variety of materials such as aluminoxanes, distancovanes, or alloxy-zinc and alloxy-aluminum compounds (see Apocatin, D.E., J.B. Lando, and J.R. Shellon, J. POLYM. Sci. PART A-1, Vol. 9, pp. 2757–2787 (1971), Gross, R.A., Y. Zhang, G. Konnad, and R.W. Lenz, MACROMOLECULES, Vol. 25, pp. 4807–4818 (1988); and Dubois, P. I. Barakat, R. Jerôme, and P. Tayssié, MACROMOLECULES, Vol. 25, pp. 4407-4412 (1993). Le Borgne, A. and N. Spassky, POLYMER, Vol. 30, pp. 2312–2319 (1988); Tanahashi, N., and Y. Dol, MACROMOLECULES, Vol. 26, pp. 4834-890 (1993); and Kennitzer, J.E., S.P. McCarthy, and R.A. Gross, MACROMOLECULES, Vol. 26, pp. 4843-890 (1993); and Kennitzer, J.E., S.P. McCarthy, and R.A. Gross, MACROMOLECULES, Vol. 26, pp. 4843-890 (1993); and Kennitzer, J.E., S.P. McCarthy, and R.A. Gross, MACROMOLECULES, Vol. 26, pp. 4843-890 (1993); and Kennitzer, J.E., S.P. McCarthy, and R.A. Gross, MACROMOLECULES, Vol. 26, pp. 4848-90 (1993); and Kennitzer, J.E., S.P. McCarthy, and R.A. Gross, MACROMOLECULES, Vol. 26, pp. 4848-90 (1993); and Kennitzer, J.E., S.P. McCarthy, and R.A. Gross, MACROMOLECULES, Vol. 26, pp. 4848-90 (1993); and Kennitzer, J.E., S.P. McCarthy, and R.A. Gross, MACROMOLECULES, Vol. 26, pp. 4848-90 (1993); and Kennitzer, Vol. 27, vol. 28, pp. 4848-90 (1993); and Kennitzer, Vol. 28, pp. 4848-90 (1993); and Kenn

[0955] The naturally derived PHAs of the present invention are isotactic and have the R absolute configuration at the stereocenters in the polymer backbone. Alternatively, isotactic polymers may be made where the configuration of the stereocenters is predominantly S. Both isotactic materiats will have the same physical properties and most of the same chemical reactivities except when a stereospecific respect, such as an enzyme, is involved. Atactic polymers, polymers with random incorporation of R and S stereocenters, can be produced from fracemic monomers and polymerization initiators or catalysts that show no preference for either enantitemer while such initiators or catalysts often Optimerze monomers of hits policial purity is isotactic polymer (s.g., distannovane catalysts) (see Hori, Y., M. Suzuki, Y. Takahashi, A. Yamaguchi, T. Nashahia, MACROMOLECULES, Vol. 59, pp. 5533–5541 (1993)). Alternatively, isotactic polymer can be produced from racemic monomers if the polymerization catalysts has an enhanced reactivity for one enantioner over

the other. Depending on the degree of preference, separate R or S stereo-homopolymers, stereo-block copolymers, or a mixture of stereo-block copolymers and stereo-homopolymers may be produced (see Le Borgne, A. and N. Spassky, N., POLYMER, Vol. 30, pp. 2312-2319 (1989); Tanahashi, N., and Y. Doi, MACROMOLECULES, Vol. 24, pp. 5732-5733 (1991); and Bervenuti, M. and R.W. Lenz, J. POLYM, SCIL: PART A: POLYM, CHEM, Vol. 29, pp. 793-805 (1991)). Some initiators or catalysts are known to produce predominantly syndrotactic polymers, polymers with alternating R and S stereocenter repeat units, from razenic monomers (see Kermitzer, J. E., S.P. McCarthy and R.A. Gross, MACRO-MOLECULES, Vol. 26, pp. 1221-1229 (1993)) while some initiators or catalysts may produce all three types of stare-polymers (see Hocking, P. J. and R.H. Marchessault, POLYM BULL., Vol. 30, pp. 163-170 (1993))

(0056) For example, preparation of poly(5-hydroxyputyrate-ao-3-hydroxyalkanoate) copolymars wherein the 3-hydroxyalkanoate comonomer is a 3-alkyl-β-propiolactone wherein the alky group contains at least three (3) carbons long, are carried out in the following manner. Proper precaulions are made to exclude air and molsture. The flactone monomers (purified, dried, and stored under inert aimosphere), β-butyrioctone and a 3-alkyl-β-propiolactone in the desired molor ratio, are charged via syringe or canula to an oven-dried, argon-progreg, and filament do brossilicate-glass tube or lask capped with a rubber septirum. The polymerization catalyst is added as a toluene solution via syringe. The tube is carefully syringed to mit the reagents (but no contact the rubber septirum) and then heated in an olb ahal that desired temperature for the prescribed time. As the reaction proceeds the mixture becomes viscous and may solidity. It isotatic polymer is produced, solid polymer precipitates out until the entire mass solidinies. The product can then be cooled, removed from the tube, and in do fresidual monomer by vacuum drying. Alternatively, the product can be dissolved in an appropriate solvent (e.g., chloroform) and recovered by precipitation in a nonsolvent (e.g., chloroform) and recovered by precipitation in anonsolvent (e.g., chloroform) solvent experiments of the standard methods such as size exclusion chromatography (SEC, also known as gel permeation chromatography or GPC). The comonomer content of the polymers is determined by nuclear magnetic resonance (NMR).

[057] In a preferred method of synthesizing the PHAs of the present invention, the initiator is an alkylzinc alkoxide, as disclosed in the US Patient No. 5,648,452 entitled "Polymerization of Beta-Substituted-Beta-Propiolactones initiated by Alkylzinc Alkoxides", L.A. Schochtman and J.J. Kemper, assigned to The Procter and Gamble Company, Issued July 1, 1997. Such initiators have the general formula R*ZnoR8, wherein R*I and R*2 are independently a C₁-C₁, alkyl. In a preferred method of synthesis, the initiator is selected from the group consisting of ethytzinc isopropoxide, methylzinc isopropoxide, or ethylzinc methoxide, or ethylzinc methoxide, more preferably ethylzinc isopropoxide.

[0058] Other copolymers useful in the present invention can be made by substituting the starting materials (monomers) in the above procedure with 3-aikyl-f-lactones corresponding to the monomer units desired in the final copolymer product. [0059] Alternalively, biological synthesis of the biodegradable PHAs useful in the present invention may be carried out by fermentation with the proper organism (natural or genetically engineered) with the proper feedstock (single or multicomponent). Biological synthesis may also be carried out with botanical species genetically engineered to express the copolymers of interest (see World Patent Application No. 93-02.187, Somewille, Polirier and Dennis, published February 4, 1993, and U.S. Patent No. 5,600.555, Dennis et al, issued July 22, 1997, and U.S. Patent No. 5,610,041, Nawmth et al., Issued March 11, 1997; and Poole, R., SCIENCE, Vol. 245, pp. 1187-1189 (1989).

Crystallinity

[0060] The volume percent crystallinity (%) of a semi-crystalline polymer (or copolymer) often determines what type of end-use properties the polymer possesses. For example, highly (greater than 50%) crystalline polyethyiene polymers are strong-and stiff, and suitable for products such as plastic milk containers. Low crystalline polyethyiene, on the other hand, is flexible and tough, and is suitable for products such as food wraps and garbage bags. Crystallinity can be determined in a number of warps, holuding x-ray diffraction, differential scanning calorimetry (DSC), density measure-

ments, and infrared absorption. The most suitable method depends upon the material being tested. [1061] X-ray diffraction is most appropriate when little is known slowth the thermal properties of the material and crystal structural changes may occur. The basic principle relies on the fact that amorphous parts of the material scatter x-rays in a diffuse of broad range go langles, white crystals diffract x-rays into sharp, precisely defined angles. The balls scattered intensity is constant, however. This allows calculation of the amount of crystaline material in a sample if the amorphous and crystalline diffracted intensities can be separated. A very precise method has been developed by Ruland, which and decid differences in percent crystallinity as small as 2% (see Vortic, C., F.J. Balta-Caleja, X-RAY SCATTERING FROM SYNTHETIC POLYMERS, Elsevier: Amsterdam, (1998); and Alexander, L., X-RAY DIFFRACTION METHODS IN POLYMERS CICNOC. New York, (1998).

[0662] Upon melting, crystals require a fixed amount of heat at the melting temperature transforming from crystalline to moiten matter. This heat of fusion can be measured by a number of thermal techniques, the most popular being DSC, if the heat of fusion of a 100% crystalline material is known, and no significant annealing, or meltirecrystallisation phenomena occur upon heating to the melt, then DSC can guite accurately determine weight fraction crystallinity (see THERMAL CHARACTERIZATION OF PCLYMER MATERIALS, E. Tuni, Ed., Academic Press, New York, (1980), and

Wunderlich, B., MACROMOLECULAR PHYSICS, Academic Press, New York, (1980)).

[0063] If the densities of the pure crystalline and pure amorphous material is known then density measurements of a material can yield the degree of crystallnity. This assumes additivity of specific volumes, but this requirement is fulfilled for polymers (or copolymers) of homogeneous structure. This method depends on careful sample preparation so that no bubbles or large voids exist in the sample.

[0064] If purely crystalline and amorphous absorption bands can be identified, then the infrared absorption spectrum offers a convenient way of determining crystallinity, (see Tadokoro, H., STRUCTURE OF CRYSTALLINE POLYMERS, John Wiley & Sons, New York, (1979)).

[0065] It should be noted that different ischrinques will often give rise to slightly different values of to, because they are based on different physical principles. For example, density measurements often give higher values than x-ray diffraction. This is due to the continuous changing of the density in the interface between crystalline and amorphous polymer (or copolymer) material. White x-ray diffraction does not detect this matter as crystalline, density measurements will be afficied by this interface region.

[0066] In general, PHAs of the present invention preferably have a crystallinity of from about 0.1% to about 99% as measured via x-ray diffraction; more preferably from about 2% to about 80%, more preferably still from about 20% to about 70%.

[0067] When a PHA of the present invention is to be processed into a film, the amount of crystallinity in such PHA is more preferably from about 2% to about 55% as measured via x-ray diffraction; more preferably from about 5% to about 50%; more preferably still from about 20% to about 40%.

20 [0068] When a PHA of the present invention is to be processed into a sheet, the amount of crystallinity in such PHA is more preferably from about 0.1% to about 50% as measured via x-ray diffraction; more preferably from about 55% to about 40%, more preferably still from about 20% to about 40%.

[0069] When a PHA of the present invention is to be processed into a normal fiber or a nonwoven fabric, the amount of crystallinity in such PHA is more preferably from about 60% to about 99% as measured via x-ray diffraction; more preferably man about 70% to about 99%, more preferably till more about 90% to both 99%, more preferably till more about 90% to both 99%.

[0070] When a PHA of the present invention is to be processed into a soft elastic fiber, the amount of crystallinity in such PHA is more preferably from about 30% to about 80% as measured via x-ray diffraction; more preferably still meabout 40% to about 80% more preferably still from about 40% to about 80% more preferably still from about 40% to about 80%.

[0071] When a PHA of the present invention is to be processed into a molded article, the amount of crystallinity in such PHA is more preferably from about 10% to about 60% as measured via x-ray diffraction; more preferably from about 30% to about 10%, more preferably sift from about 30% to about 60%.

[0072] When a PHA of the present invention is to be processed into an elastomer or adhesive, the amount of crystallinity in such PHA is more preferably less than about 50% as measured via x-ray diffraction; more preferably sell less than about 50%, more preferably still less than about 50%, more preferably still less than about 50%, more preferably still less than about 50%.

Melt Temperature

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[0073] Preferably, the biodegradable PHAs of the present invention have a melt temperature (Tm) of from about 30°C to about 160°C, more preferably from about 60°C to about 140°C, more preferably still from about 90°C to about 120°C.

Plastic Articles Comprising PHA

[0074] The PHAs of the present invention can be processed into a variety of plastic articles, including but not limited to sheets, fibers, foams, molded articles, nonwoven fabrics, elastomers, and adhesives.

[0075] Prior to Applicants' invention, polyhydroxyalkanoates studied for use in commercial plastics production presented significant impediments to their use in plastics. As discussed above, polyhydroxyalkanoates such as PHB and the copolymer PHBV are difficult to process due to their thermal instability, Applicants surprisingly discovered, such linear, random copolymers with a limited number of medium sized branched alkyl chains containing three (3) carbons. , provide, in addition to biologradobility, the following properties, particularly as compared to PHB or PHBV: a) a lower melt temperature, b) a lower degree of crystallinity, and c) an improved melt rheology. This is especially surprising in light of the fact that the longest straight branch of the medium sized branched alkyl chain contains only two (2) carbons. [0076] Without being bound by theory, Applicants betieve characteristics a) and b) are achieved by exclusion of the second RRMU from the crystal lattice of the first RRMU, thereby resulting in a decreased temperature for thematic characteristic c) is achieved by increased enlanglement between the coopwher chains due to the side chains of the second RRMU. Such increased enlanglement any occur by increased hydrodynamic volume of the copolymer (e.g., the second monomer unit creates kinks in the helical structure), the Thooking" or "calching" of the side chains of different copolymer backbones white in the melt, or the decreased chains sistend use to the work Tm (i.e., the enlarged thermal process window).

B. Sheets

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[0077] In another embodiment of the present invention, the plastic article is a sheet. As used herein, "sheet" means a very thin continuous piece of a substance, having a high length to thickness ratio and a high width to thickness ratio, wherein the material is thicken than 0.254 mm. Sheeting shares many of the same characteristics as film in terms of properties and manufacture, with the exception that sheeting is stiffer, and has a self-supporting nature. Such differences in stiffness and support result in some modification of the manufacturing methods.

1, Methods of Manufacture

[0078] Sheets, because of thickness and consequent stiffness, cannot be blown as a film. However many other of the same processes used to make film can be modified to make sheeting. One example is cast extrusion which is described previously. In addition to extrusion, sheeting is also made via rolling and calendering.

a. Rolling

[0079] Rolling produces a film with predominately machine direction orientation by accelerating the film from a nip point where the thickness is reduced (ENCYCLOPEDIA OF POLYMER SCIENCE AND ENGINEERING, Vol. 8, pp. 88-106, John Wiley and Sons, New York, (1986); hereinafter referred to as "EPSE4"). Large forces are lound at the nip point, but overall orientation can be increased over other forms of machine direction orientation.

b. Calendering

[0080] To produce an unoriented cast film or sheet with high throughput, calendering is used (G. W. Eghmy, Jr. in MODERN PLASTICS, J. Agrandoff, ed. Encyclopedia, Vol 59(10A), pp. 220-22(1982) and R. A. Elden and A. D. Swan, CALENDERING OF PLASTICS, American Elsevier Co., Inc., New York, (1971). The calendering process employs stacks of specially hardened, driven rolls, supported in a manner so they may be bent or skewed in position relative to each other during operation. This is to control thickness in the calenderder material. Calenders are usually made up of our rolls that form three nips. These nips are the feed, metering and finishing nips. The feed nip is where the polymer is supplied, mixed, and heated. The metering nip reduces the thickness of the sheet to the approximate final thickness. The finishing nia adults the gauge of the sheet by varying the position of the third or middle roll. (see EPSE-2)

35 C. Fibers

[0841] In another embodiment of the present invention, the plastic article is a fiber. As used herein, "fiber frefers to a flexible, macroscopically homogeneous body having a high length-howidth ratio and a small cross section. A general eventive of fiber scan be found in the ENCYCLOPEDIA OF POLYMER SCIENCE AND ENGINEERING, Vol. 6, p. 647-755 and pp. 802-839, John Wiley and Sors, New York, (1986) fiverlenather referred to as "EPSE-2"). The fibers of the present inventions are useful as textiles in years for garments. The fibers of the present invention are also useful for manufacturing lightweight fibrous materials useful in agricultural applications to protect, promote, or control plant growth. They are also used in green house thermal screenes, corp row covers, furl dozers, weed barriers and hydroponics. Key properties are light, air, and moisture permeability. An important aspect is cost effectiveness when considered in terms of weight, strength, and dimension stability.

[0882] An elastomeric fiber is a fiber that consists of polymers (or copolymers) with a main glass transition temperature much below orom temperature (see EPSE-2). This criterion excludes some fibers with elastic properties, such as crimped hard fibers, but allows inclusion for multi-consistuant fibers where one of the constituents is an elastomeric fiber are characterized by a higher elongation at break, lower modulus, and higher recovery from large deformation than normal fibers.

1. Methods of Fiber Manufacture

[0083] The fibers of the present invention may be processed using a variety of conventional techniques well-known in the art including, but not limited to, mell spinning, dry spinning, and wet spinning. Combinations of these three basic processes are often used.

[0084] In melt spinning a PHA of the present invention is heated above its melting point and the molten PHA is forced through a spinneret. A spinneret is a die with many small orifices which are varied in number, shape and diameter (see

EPSE-2). The jet of molten PHA is passed through a cooling zone where the PHA solidifies and is then transferred to post-drawing and take-up equipment.

[0085] In dry spinning, a PHA of the present invention is dissolved and the PHA solution is extruded under pressure through a spinneret (see EPSE-2). The jet of PHA solution is passed through a healting zone where the solvent evaporates and the filament solidifies.

[086] In wet spinning, a PHA of the present invention is also dissolved and the solution is forced through a spinneret which is submerged in a coagulation bath (see ESPE-2). As the PHA solution emerges from the spinneret onfices within the coagulation bath, the PHA is either precipitated or chemically regenerated. Usually, all these processes need further drawing for useful properties to be obtained, for example to serve as textell fibers. Drawing refers to stretching and attenuation of fibers to achieve an inverestible extension, induce moticair ententation, and develop a fiber-line structure (see ESPE-2). This fine structure is characterized by a high degree of crystallinity and by orientation of both the crystallites and the amorphous PHA chain segments.

D. Foams

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[0087] In another embodiment of the present invention, the plastic article is a flexible foam. As used herein, "foam" refers PHA of the present invention whose apparent density has been substantially decreased by the presence of numerous cells distributed throughout its bulk (see ASTM D 883-82T, American Society for Testing and Maleriats, Philadelphia, Pa. (1962)). Such two-phase gas/solid systems in which the solid is continuous and composed of a synthetic polymer or nubber include cellular polymers (or copolymens), expanded plastics and foamed plastics (ENCY-CLOPEDIA OF CHEMICAL TECHNOLOGY, Vol. 11, John Wiley & Sons, New York (1980), hereinafter referred to as a recommendation of the plastic of the plastic in the plastic interval.

[0088] The gas phase is distributed into pockets or voids called cells which are classified into two types, open and closed. Open-celled material are foams whose cells are inter-connected such that gases may pass freely through the cells. Closed-cell materials have cells that are discrete and isolated from each other.

Soms are further categorized into flexible and rigid Gams. This classification is based on a particular ASTM

test procedure (see ASTM D, Vol. 37, pp. 1566-1578, American Society of Testing and Materials, Philadelphia, Pa. (1978)), A flexible foam is a foam which does not rupture when a 20 x 2.5 x 2.5 or piece is wrapped around a 2.5 cm mandrel at a uniform rate of 1 lap5s at 15-25°C, Foams that do rupture under this test are referred to as rigid barns. [0080] Foams find many applications as packaging, comfort cushioning, insulation, and structural components. In the some areas of packaging a foamed material having increased biodeyr adability and/or compostability would offer superior benefits to currently used packaging such as polystyrene, paper and starch foams. In hot food containers, polystyrene offers significantly higher thermal insulation over the only currently degradable alternative, paper wraps. Foamed articles comprising a PHA of the present invention have the thermal insulating properties of polystyrene, yet are biodegradable

andior compostable. These materials are ideal for hot food take-out and cold food packaging.

(0091) Foamed polystyrene chips are used as cushioned packing materials for consumer and industrial goods. Many
of these chips end up in landfills. Foamed chips comprising a PHA of the present invention perform as well as polystyrene
and have increased biodegradability and/or compostability. Unlike other compostable packaging material such as starch,
such PHA chips are resistant to many common solvents and liquids including water.

1, Methods of Foam Manufacture

[0092] The foams of the present invention may be processed using conventional procedures well-known to those skilled in the art. A predominant method of foam production involves expanding a fluid polymer (or copolymer) phase to a low density cellular phase and then preserving this state (see ECT). Other processes include leaching out materials that have been previously dispersed in the polymer (or copolymer), sintering small particles and dispersing cellular particles in a polymer (or copolymer). Three sites make up the expansion process. These are cell initiation, cell growth and cell stabilization. Many methods are used to create, grow, and stabilize cells.

[0093] Expandable formulations rely on increasing the pressure within the initiated cells relative to that of the surroundings. The cells are stabilized by either chemical (e.g. crosslinking, polymerization) or physical means (crystallization) multi-glass transition). Polystyrene is an example of a polymer that is foamed by of this kind of process. A blowing agent such as isomeric pentanes and hexanes or halocarbons (H. R. Lasman, MODERN PLASTICS, Vol. 42(1A), p. 314 (1964)) is mixed with the polymer (or copplymer) either by heating and allowing the bolving agent to pentarte the polymer (U.S. Pat. 2,861,321, issued June 15, 1964, F. Stasthy and R. Gaeth, assigned to BASF), or by polymerizing the polystyrene in the presence of the blowing agent (U.S. Pat. 2,863,802, issuel May 9,1961, G. F. D'Alelio, assigned to Koppers Co.). Fabrication of articles are usually carried out in multiple steps, the first of which uses steam, hot water or hot air to expand the polymer into low density preformed beads. These preformed beads are aged, sometimes in multiple steps for correct cell size, and then packed into motis and fused together by heat and further expansion (S. J.).

Skinner, S. Baxter, and P.J. Grey, Trans. J. PLAST, INST. Vol. 32, p. 180 (1964)). Stabilization is accomplished by cooling the polymer to temperatures below its glass transition temperature.

[0094] Decompression expansion processes create and grow cells by lowering the external pressure during processing. Cellular polyethylene and polyproplene are often made in this manner. A decomposing blowing agent is premixed with the polymer for oppolymer and fed through an extruder under elevated temperature and pressure such that the blowing agent partially decomposes. When the material exits the extruder, it enters a lower pressure zone. Simultaneous expansion and cooling take place, resulting in a stable cellular structure owing to rapid crystallization of the polymer (R. H, Hansen, SPE J., VJ. 18, p. 7 (1962), W. T. Higgins, M.O.P. LAST., Vol. 31 (7), p. 99, (1954).

[0055] Dispersion processes produce fearns by directing dispersing solld or gas into the polymer (or copolymer) phase and then, when necessary, stabilizing the mixture (ECT). In one such process, frothing, a gas is mechanically dispersed in the polymer or monomer phase, producing a fearn of temporary stability. This Goaria is then chemically stabilized by crosslinking or polymerazion. Latex fearn rubber is manufactured in this way (see ECT).

E. Molded Articles

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[0096] In another embodiment of the present invention, the plastic article is a molded article. As used herein, "molded article" means objects that are formed from polymer or copolymer materials (e.g., PHA) which are injected, compressed, or blown by means of a gas into shape defined by a female mold. These objects can be solid objects like toys, or hollow like bottles and containers.

[0037] Injection moding of thermoplastics is a multi-step process by which a PHA of the present invention is heated until it is moiten, then forced into a closed mold where it is shaped, and finally solidied by cooling. There are a variety of machines that are used in injection molding. Three common types are ram, screw plasticator with injection, and reciprocating screw devices (see ENCY-COPEDIA OF POLYMER SCIENCE AND ENGINEERING, 49.8, pp. 102-138). John Wiley and Sons, New York, (1986); hereintaire referred to as "EPSE-37. A ram injection molding machine is composed of a cylinder, spreader, and plunger. The plunger forces the melt in the mold. A screw plasticator with a second stage injection consists of a plasticator, directional valve, a cylinder without a spreader, and a ram. After plastication by the screw, the screw rotates to melt and mit the market and then moves forward to frore the melt into the mold. [0048]

Compression molding in thermoplastics consists of charging a quantity of a PHA of the present invention in hellower half of an open die. The open add between where other brought together under pressure, and them mottees.

PHA conforms to the shape of the die. The mold is then cooled to harden the plastic (see EPSE-3).

[0099] Blow molding is used for producing bottles and other hollow objects (see EPSE-3). In this process, a tube of
molten PHA known as a parison is extruded into a dosed, hollow mold. The parison is then expanded by a gas, thrusting
the PHA against the walls of a mold. Subsequent cooling hardens the plastic. The mold is then opened and the article
semonal.

[0100] Blow molding has a number of advantages over injection molding. The pressures used are much lower than injection molding. Blow molding can be typically accomplished at pressures of 25-100 psi between the plastic and the mold surface. By comparison, injection molding pressures can reach 10,000 to 20,000 psi (see EPSE-3). In cases where the PHA has a have molecular weights too high for easy flow through molds, blow molding is the technique of choice. High molecular weight polymers (or copolymers) often have better properties than low molecular weight analogs, for example high molecular weight malerals have greater resistance to environmental stress cracking. (see EPSE-3). It is possible to make extremely thin walls in products with blow molding. This means less PHA is used, and solidification times are shorter, resulting in lower costs through material conservation and higher throughput. Another important feature of blow molding is that since it uses only a temale mold, slight changes in extrusion conditions at the parison nozzic can vary well binkness (see EPSE-3). This is an advantage with siructures whose necessary wall thicknesses cannot be predicted in advance. Evaluation of articles of several thicknesses can be undertaken, and the thirmest, thus lightest and chappets, article that meets specifications can be used.

F. Nonwovens

[0101] In another embodiment of the present invention, the plastic article is a nonwoven. As used herein "nomoven" means porous, textile like materials, usually in flat sheet form, composed primarily, or entirely, of fibers assembled in webs that are manufactured by processes other than spinning, weaving, or knitting. A general overview of nomovore flatincs can be found in the ENCYCLOPEDIA OF POLYMER SCIENCE AND ENGINEERING, Second Edition, Vol. 10, pp. 204-226 (referred to hereafter as "EPSE-4"). Other names for these materials are bonded fabrics, formed fabrics, or engineered fabrics. The thickness of the fabric sheets may vary from 25 mm to several centimeters, and the weight from 10 gim² to 1 kg/m². Nomoven fabrics have a wide range of physical properties depending on the material and process used in forming the week. A fabric may be self-supporting and stiff as paper or drapable as a conventional cloth.

fabric

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[0102] In contrast to conventional textiles, the fundamental structure of all nonwovens is a web of fibers arranged more or less randomly (NONWOVENS IND., Vol. 17, p. 36 (Mar. 1986), NONWOVENS WORLD, Vol. 1, p. 36 (May-June 1986)). Thus, the key element is the single fiber. Tensile, tear, and tactile properties in the nonwoven arise from adhesive or other chemical and physical bonding, fiber-to-fiber friction created by entanglement, and reinforcement by other materials such as foams and films (see EPSE-4).

Method of Manufacture of Nonwoven Fabrics

[0103] The nonwoven fabrics of the present invention may be made by conventional techniques known in the art. The production of nonwoven fabrics involves: 1) making fibers of various lengths and diameters; 2) creating a web of these fibers; and 3) bonding of fibers within the web by adhesive, or mechanical-frictional forces created by fiber contact or entanglement. In addition to these steps, reinforcing the web by forming a composite with other materials (e.g., yarns, scrims, films, nettings, and unbonded webs) is sometimes preferred. Variations of one or several of these steps allows for the enormous range of nonwoven fiber types. The term "staple fibers" was originally applied to fibers of natural origin long enough to be processed on textile machinery, but excluding endless filaments, eg, silk. In the present context, as applied to PHA of the present invention, "staple fibers" are of relatively uniform length, ca. 1.3-10.2 cm, with a regular crimp i.e., a three-dimensional wavelike shape. Regenerated and other extruded fibers are endless as formed. They are cut during the manufacturing process to a specified length to meet a processing or market need. Extruded fibers are also produced as continuous filaments without crimp. The processes for forming webs from staple fibers are different from those using continuous filaments. The products obtained from staple and filament fiber webs may differ substantially in properties (see EPSE-4).

[0104] The mechanical properties of the fibers as defined by their chemical composition, determine the ultimate properties of the fabric. Web structure and bonding maximize inherent fiber characteristics (see EPSE-4). Other materials that may be used in the nonwovens of the present invention in combination with the PHA are wood pulp; regenerated fibers including viscose rayon and cellulose acetate; and synthetic fibers like poly(ethylene terephthalate) (PET), nylon-6, nylon 6,6, polypropylene (PP), and poly(vinyl alcohol). Facings of disposable diapers or sanitary napkins made from PHA nonwoven fabrics of the present invention preferably feel dry even when the absorbent, inner absorbent layer is saturated. Important fiber characteristics that affect performance include length, diameter, density, crimp, cross section shape, spin-finish (lubricant that is added to the surface of extruded fibers to enhance processability), delustering (small amounts of TiO₂ pigment added before extrusion to increase whiteness or to reduce sheen) and the draw ratio.

a. Web-making methods

[0105] The characteristics of the fiber web determine the physical properties of the final product. These characteristics depend largely on fiber architecture, which is determined by the mode of web formation. Fiber architecture includes the predominant fiber direction, whether oriented or random, fiber shape (straight, hooked, or curled), the extent of interfiber engagement or entanglement, crimp, and compaction (web-density control). Web characteristics are also influenced by fiber diameter, length, web weight, and chemical and mechanical properties of the polymer (see EPSE-4).

[0106] The choice of method for forming the web is determined by fiber length. Initially, the methods for forming webs from staple-length fibers (fibers long enough to be handled by conventional spinning equipment, usually from about 1.2 to about 20 cm long, but not endless) are based on the textile-carding process, whereas web formation from short fibers is based on papermaking technologies. Although these technologies are still in use, other methods have been subsequently developed. For example, webs are formed from long, virtually endless filaments directly from bulk polymer; both web and fibers are produced simultaneously (see EPSE-4). A variety of web-making methods are known, including carding, air-laying, wet-forming, spinbonding, and meltblowing.

[0107] The carding process is derived from the ancient manual methods of fiber carding, where natural staple fibers were manipulated by beds of needles. In carding, clumps of staple fibers are separated mechanically into individual fibers and formed into a coherent web by the mechanical action of moving beds of closely spaced needles.

[0108] In the air-laying process, the orientation created by carding is effectively improved by capturing fibers on a screen from an airstream (see US Palent No. 3,338,992, G. A. Kinney, assigned to E.I. du Pont de Nemours & Co., Inc., issued August 29, 1967). The fibers are separated by teeth or needles and introduced into an airstream. Total randomization would exclude any preferential orientation when the fibers are collected on the screen.

[0109] Wet-forming processes employ very short fibers. Initially, webs are formed from short fibers by modified papermaking techniques. The fibers are continuously dispersed in a large volume of water and caught on a moving endless wire screen. Once the web is caught on the screen, it is transferred to belts or felts and dried on heated drums (see EPSE-4).

[0110] The spunbonded web process involves making fibers and web simultaneously, directly from bulk polymer. The

bulk polymer is melted, extruded, and drawn (often by theolectaric forces) to filaments that are randomized and deposited onto belts as a continuous web. The filaments are virtually endless. The spunhout process produces webs of low crimp filaments in the normal diameter range of about 1.7 dex (1.5 den) or signity higher. The birefringence and uniform of diameter of these filaments are similar to standard textile interns and filaments (see EPSE-4). Each production line) is suitable for a specific polymer and a single-bonding system (see U.S. Pat. 4, 163.305 (Aug. 7, 1979), V. Semjonow and

J. Foedrowitz (to Hoechst AG)).
[0111] Webs are also made directly from bulk polymers by the meltblown process (see US Patent No. 3,322,607, S.L. Jung, assigned to E.I. duPont de Nemours & Co., Inc., May 30, 1967). The molten PHA is forced through very fine holes in a special die into a high velocity airstream where the PHA is formed into very fine, athough irregular, filaments of indeterminate lengths. The filaments are simultaneously formed into a web where meiting and resolidification, and prossibly static forces, hold them together (see EPSE-d). The web consists primarily of filaments with very fine diameters.

b. Web bonding

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[0112] The bonding of fibers gives the strength to the web and influences other properties. Both adhesive and mechanical means are used. Machanical bonding emolys he engagement of fibers by frictional forces. Bonding can also be achieved by chemical reaction, i.e., formation of covalent bonds between binder and fibers (see EPSE-4).

G. Elastomers

[0113] In another embodiment of the present invention, the plastic article is an elastomer. As used herein "elastomer' refers to materials which schibit both long-range deformability on application of stress and essentially complete recovery on removal. A general discussion on elastomers can be found in the Encyclopedia of Polymer Science and Engineering. Second Edition, Vol. 5, pp. 106-127 (hereafter referred to as "EPSE-57). Preferably, an elastomer of the present invention, at room temperature, can be stretched repeatedly to at least twice its original length and, after removal of the tensite olds, will immediately and forcibly return to approximately its original length. Elastomers of the present invention are above the glass-transition temperature Tg and amorphous in the unstressed state to exhibit high local segmental immolitiey necessary for deformation. The chains are floxible and intermolecular (interchain) forces are weak. The elastomers of the present invention possess a sufficient number of chemical or physical cross-links to form a continuous network in

order to restrain chain slippage.

[0114] Thermoplastic elastomers of the present invention have many of the properties of conventional elastomers such as vulcanized rubbers, but are processed as thermoplastics rather than thermosets. Transition from a fluid melt to a solid is reversible. Thermoplastic elastomers of the present invention are multiphase systems, where at least one phase is soft and rubbery and another hard. With thermoplastic elastomers, the transition from a processible melt to a solid, rubberliftic object is rapid and reversible and takes place upon cooling. Preferably, PHAs of the present invention which are processed into an elastomer have sufficiently high branch content to enable them to act as thermoplastic elastomers, with the crystalline areas acting as the hard segment and the amorphous segments acting as the soft segment. Thermoplastic elastomers of the present invention can be processed on conventional plastics equipment, such as injection molders.

40 (2015) Important structural parameters for thermoplastic elastomers are the molecular weight, the nature of the soft and hard segments, and the ratio of soft to hard segments. The ratio of hard to soft segments effects the total modulus of the elastomer, increasing with the proportion of the hard segments.

[0116] Elastomers of the present invention comprising a PHA of the present invention can also be used in blend formulations withother polymers (or copolymers), even non-distoment PHAs, to increase impact strength and toughness in stiffer malerails.

H. Adhesive

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[0117] in another embodiment of the present invention, the plastic article is an adhesive. As used herein "adhesive" means a material that joins two other materials, called adherends, together, & general discussion on adhesives can be found in the Encyclopedia of polymer Science and Engineering, Vol. 1, pp. 547-577, (hereafter referred to as "EPSE-6"), In one embodiment of the present invention, the adhesive is applied as a fliquid, preferably of a low viscosity, In the fliquid form the adhesive west between adherend surfaces. The liquid form of the adhesive wis obtained by heating to the point that flow occurs, dissolving or dispersing the material in a solvent, or starting with liquid monomers or oligomers that polymerize or react after application. The adhesive then undergoes a phase change to a solid either by cooling, solvent evaporation, or reaction, in order for the joint to acquire the necessary strength to resist shearing forces. However, pressure-sensitive adhesives are an exception, since no phase change occurs.

[0118] The PHAs of the present invention may be processed into a variety of adhesives, including but not limited to, not melt, solution, dispersion and pressure sensitive adhesives.

1, Hot-melt Adhesives.

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[0119] As used herein, "hot-mail adhesive" refers to a themoplestic polymer or copolymer (e.g., a PHA of the present invention) that is heated to obtain a liquid of flowable viscosity, and, after application, cooled to obtain a solid. Generally, the molecular weight of the adhesive is fatlored to provide flowability in the melt, but still be strong enough in the solid form to resist shearing forces experienced in the application. Due to their themoplestic properties, the PHAs of the present invention are particularly useful as hot-mail adhesives. The primary feature of hot-melt adhesives is the ability of the thermoplestic material to flow above a certain temperature, high above the normal use temperature of the bond. Upon cooling, the material handless, either through passing through the glass transition temperature of one of the components, or the crystallization temperature. This hardening lends physical integrity to the bond. In PHAs, the mode of solidification is crystallization.

Solutions and dispersions.

[0120] The adhesives of the present invention may be applied either as solutions, in water or an organic solvent, or in the form of squeous dispersions. In either form, the solvent must be removed after application for the adhesive to attain the required solid form. The solution or dispersion is usually applied to one of the surfaces to be bonded, and the solvent removed before the second surface is joined; often, healing is required to expedite the drying step. With porous substrates, such as paper or wood, final drying can take place after formation of the joint. Solids contents of the solutions vary from \$5 to \$9%, although values from 20 to 50% are most common.

[0121] As used herein, "dispersion" refers to when adhesives are prepared by true emulsion polymerization or dispersed as larger particles in some carrier fluid. In addition to their economic advantage, dispersions containing d45-50% solids offer lower viscosity than solutions, even if the solids are high molecutar-weight polymers (EPSE-6). Adhesive dispersions of the present invention may be prepared by high shear in the presence of surfactants to obtain waterborne formulations, procedures which are well known to those skilled in the art.

3, Pressure-sensitive Adhesives.

[0122] Another type of adhesive of the present invention is a pressure-sensitive adhesive. Unlike other adhesives, the pressure-sensitive adhesives do not change their physical state from the initial application, to the final breaking of the adhesive bond. They emain permanently deformable, and may after under even slight application of pressure. They are adhesives that in dry form are permanently tacky at room temperature and that firmly adhere to surfaces upon mere contact. The most common form of pressure-sensitive adhesive is on a backing, usually in tape form. Common masking tape, for example, is manually applied after the user removes the desired length from a roll. Many bandages are held to the skin by pressure-sensitive adhesives.

40 Disposable Personal Care Products

[0123] The present invention further relates to disposable personal care products comprising a PHA of the present invention. For example, compostable absorbent articles comprising a liquid previous topsheet, a liquid impervious back-steet comprising a film of the present invention (i.e., a film comprising a PHA of the present invention), and an absorbent core positioned between the topsheet and backsheet. Such absorbent articles include infant diapers, adult incontinent herists and nads, and fermine hydrice pads and literes.

[0124] Additional personal care products comprising a PHA of the present invention include personal cleansing wipes; disposable health care products such as bandages, wound dressings, wound cleansing pads, surgical gowns, surgical covers, surgical pads; other institutional and health care disposables such as gowns, wipes, pads, bedding items such as sheets and pillowcases, foam mattress pads.

A. Absorbent Articles

[0125] Films of the present invention used as liquid imporvious backsheets in absorbent articles of the present invention, such as disposable diapers, typically have a thickness of from 0.01 mm to about 0.2 mm, preferably from 0.012 mm to about 0.2 mm, preferably from 0.012 mm to

[0126] In general, the liquid impervious backsheet is combined with a liquid pervious topsheet and an absorbent core positioned between the topsheet and the backsheet. Optionally, elastic members and tape tab fasteners can be included.

- While the topsheet, the backsheet, the absorbent core and elastic members may be assembled in a variety of well known configurations, a preferred disper configuration is described generally in U.S. Patent 3,860,003, entitled "Contracible Side Portion for bisposable Diagnet" which issued to Kenneth B. Buell on January 14, 1975.
- [0127] The topsheet is preferably, solt-feeling, and non-irritating to the wearer's skin. Further, the topsheet is liquid pervious, permitting liquids to readily penetrate through its thickness. A suitable topsheet may be manufactured from a wide range of materials such as porous foams, reliculated foams, paertured plassic films, natural fibers (e.g., wood or cotton fibers), synthetic fibers (e.g., polyester or polypropylene fibers) or from a combination of natural and synthetic fibers. Preferably, the topsheet is made of a hydrophobic material to solate the wearer's skin from liquids in the absorbent core.
- 10 [0128] A particularly preferred topsheet comprises staple-length fibers having a denier of about 1.5. As used herein, the term "staple-length fibers" refers to those fibers having a length of at least about 16 mm.
 - [123] There are a number of manufacturing techniques which may be used to manufacture the topsheet. For example, the topsheet may be woven, non-woven, spunbonded, carded, or the like. A preferred topsheet is carded, and thermally bonded by means well known to hose skilled in the fabrics art. Preferably, the topsheet has a weight from about 18 to about 25 g/m², a minimum dried tensile strength of at least about 400 g/cm in the machine direction, and a wet tensile strength of a least about 50 g/m² in the ross-machine direction.
 - [0130] In a preferred embodiment of the present invention, the top sheet comprises a PHA of the present invention.

 [0131] The topsheet and the backsheet are joined together in any suitable manner. As used herein, the term 'joined' encompasses configurations whereby the topsheet is directly joined to the backsheet by affixing the topsheet directly to the backsheet, and configurations whereby the lopsheet is indirectly joined to the backsheet by affixing the topsheet of the backsheet and configurations whereby the lopsheet is indirectly joined to the backsheet and to configurations whereby the lopsheet is indirectly joined to the backsheet and to configurations whereby the lopsheet is indirectly joined to the backsheet and to configurations whereby the lopsheet is indirectly joined to the backsheet and to configurations whereby the lopsheet is indirectly joined to the backsheet and the lopsheet in the lopsheet in any lopsheet and the lopsheet in the lopsheet in
- 20 to the backsheel, and configurations whereby the topsheet is indirectly joined to the backsheet by affixing the topsheet to intermediate members which in turn are affect of the backsheet. In a preferred embodiment, the topsheet and the backsheet are affixed directly to each other in the diaper periphery by attachment means such as an adhesive or any other attachment means known in the art. For example, a uniform, continuous layer of adhesive, a patterned layer of adhesive, or an array of separate lines or spots of adhesive may be used to affix the topsheet to the backsheet.
- 25 [0132] In a preferred embodiment of the present invention, the adhesive comprises a PHA of the present invention [0133]. Tape tab its steners are typically applied to the back waisthand region of the diaper to provide a fastening mass for holding the diaper on the wearer. The tape tab fasteners can be any of those well known in the art, such as the fastening tape disclosed in U.S. Patent 3,948,594 issued to Kenneth B. Buell on November 19, 1974. These tape tab fasteners or other diaper fastening means are typically applied near the corners of the diaper.
- 30 [0134] Preferred dispers have elastic members disposed adjacent the periphery of the disper, preferably along each longitudinal edge so that the elastic members and to draw and hoth the disper against the legs of the waver. The legals members are secured to the disper in an contractible condition so that in a normally unrestrained configuration the elastic members effectively contract or gather the disper. The elastic members can be secured in an contractible condition at least two ways. For example, the elastic members may be stretched and secured while the disper is in an uncontracted condition. Alternatively, the disper may be contracted, for example; by pleasing, an elastic member secured and connected to the disper while the elastic members are in their relaxed or unstretched condition.
 - [0135] The elastic members may take a multihude of configurations. For example, the width of the elastic members may be varied from about 0.25 mm or about 25 mm or more; the elastic members may comprise a single stand of elastic members may be retargular or curvilinear. Still further, the elastic members may be efficied to the diaper in any of several ways which are known in the art. For example the elastic members may be ultrasonically bonded, heat and pressure sealed into the diaper valing a variety of bonding patterns, or the elastic members may simply bonded, heat and pressure sealed into the diaper valing a variety of bonding patterns, or the elastic members may simply the provided in the diaper valing a variety of bonding patterns, or the elastic members may simply the provided in the diaper valing a variety of bonding patterns, or the elastic members may simply the provided between the provided patterns and the provided patterns are considered to the diaper valing the provided patterns are considered to the diaper valing the provided patterns are considered to the diaper valing the provided patterns are considered to the diaper valing the patterns are considered to the diaper valing the provided patterns are considered to the diaper valing the provided patterns are considered to the diaper valing the provided patterns are considered to the diaper valing the patterns are considered to the diaper valing the provided patterns are considered to the diaper valing the patterns are
 - be glued to the diaper.

 [0138] In a preferred embodiment of the present invention, the etastic members comprise a PHA of the present invention.
- [0137] The absorbent core of the diaper is positioned between the topsheet and backsheet. The absorbent core may be manufactured in a wide variety of sizes and shapes (e.g., rectangular, hour-glass, asymmetrical, etc.) and from a wide variety of materials. The total absorbent capacity of the absorbent core should, however, be compatible with the designed liquid loading for the intended use of the absorbent article or disper. Further, the size and absorbent capacity of the absorbent core may vary to accommodate wearers ranging from intents through adults.
- (0 138) A preferred embodiment of the diaper has an hour-glass shaped absorbent core. The absorbent core is preferably an absorbent member comprising a web or batt of airfelt, wood pulp fibers, and/or a particulate absorbent polymeric composition disposed therein.
 - [0139] In a preferred embodiment of the present invention, the absorbent polymeric composition of the absorbent core comprises a PHA of the present invention.
- 55 (0140) Other examples of absorbent articles according to the present invention are sanitary napkins designed to receive and contain vaginal discharges such as menses. Disposable sanitary napkins are designed to be held adjocent to the human body through the agency of a garment, such as an undergament or a panty or by a specially deadbett. Examples of the kinds of sanitary napkins to which the present invention is readily adapted are shown in U.S. Patent

4,697,478, entitled "Shaped Sanitary Napkin With Flaps" which issued to Kees J. Van Tiburg on August 18, 1987, and in U.S. Pakert 4,599,876, entitled "Sanitary Napkin" which issued to Kees J. Van Tiburg on May 20, 1986. It will be apparent that the films of the present invention one-prising a PHA of the present invention described herain may be used as the liquid impervious backsheet of such sanitary napkins. On the other hand it will be understood the present invention is not limited to any specific sanitary napkin configuration or structure.

[0141] In general, sanitary napkins comprise a liquid impervious backsheet, a liquid pervious topsheet, and an absorbent core placed between the backsheet and the topsheet. The backsheet comprises a PHA of the present invention. The topsheet may comprise any of the topsheet materials discussed with respect to diagres. The adhesives used in may also comprise a PHA of the present invention. The absorbent core may comprise any of the absorbent core materials.

discussed with respect to diapers, including a PHA of the present invention.

[0142] Importantly, the absorbent articles according to the present invention are biodegradable and/or compostable to a greater extent than conventional absorbent articles which employ materials such as a polyolefin (e.g., a polyethylene) harksheel.

FXAMPLE 1

Poly(3-hydroxybutyrate-co-3-hydroxy-4-methylvalerate)

[0143] Poly(3-hydrox)butyrate-co-3-hydroxy-4-methywiserate) is prepared according to the general methods described above and based on the published procedure of Hon et al. (Hon, Y., M. Suzuki, Y. Takahashia, A. Yomagushi, and T. Nishishita, MACROMOLECULES, Vol. 26, pp. 5533-5534 (1993)) for the polymertzation of [b-butyrolacione, Specifically, purified [S]-3-methylpropiolacione ([S]-B-butyrolacione), [S]-3-goryolacione, and [S]-3-boryolypropiolacione) (S0.30). In the month of the second of the second

35 FXAMPLE 2

Poly(3-hydroxyvalerate-co-3-hydroxy-4-methylvalerate)

[0144] Poly(3-hydroxyvalerate-co-3-hydroxy-4-methylvalerate) is prepared by following the same procedure as in 0 Example 1, with the exception that [§3-a-thylpropiolactone (9.50 g, 94.9 mmd) and [§]-3-topropylpropiolactone (0.71 g, 5.0 mmd) are used as the monomer charge.

EXAMPLE 3

45 Poly(3-hydroxybutyrate-co-3-hydroxyvalerate-co-3-hydroxy-4-methylvalerate)

[0145] Poly(3-hydroxybutyrate-co-3-hydroxyvaterate-co-3-hydroxy-4-methylvaterate) is prepared by following the same procodure as in Example 1, with the exception that [Is3-methylpropioiscober (7.2 o.g. 8.3.6 mmol), [Is3-sethylpropiolactone (1.7.4 g. 1.14 mmol), and [Is3-setyonylpropiolactone (1.7.4 g. 1.04 mmol) are used as the monomer charge.

EXAMPLE 4

Poly(3-hydroxybutyrate-co-3-hydroxy-4-methylvalerate-co-3-hydroxyoctanoate)

[0146] Poly(3-hydroxybulyrate-co-3-hydroxy-4-methylvalerate-co-3-hydroxyoctanoate) is propased by following the same procedure as in Example 1, with the exception that [S]-3-methylpropications (9.50 g, 110 mmol), [S]-3-isoproplypropication (o.41 g, 2.9 mmol), and [S]-3-polypropications (0.50 g, 2.9 mmol) are used as the monomer charge.

EXAMPLE 5

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Poly(3-hydroxybutyrate-co-3-hydroxyvalerate-co-3-hydroxy-4-methylvalerate-co-3-hydroxyoctanoate)

- 5 [0147] Poly(3-hydroxybutyrate-co-3-hydroxyvaterate-co-3-hydroxy-4-methy/vaterate-co-3-hydroxyoctanoate) is prepared by following the same procedure as in Example 1, with the exception that [S}-3-methy/propiolactone (7.20 g. 83.6 mmol), [S]-3-ethy/propiolactone (1.14 g. 11.4 mmol), [S]-3-isopropy/propiolactone (0.36 g. 2.5 mmol) are used as the monomer charge.
- 10 EXAMPLE 6 not example of the invention

Compostable Single Layer Film

[0148] Poly(3-hydroxybutyrate-co-3-hydroxy-4-methylvalerate) copolymer (PHBMV) of composition 5 moles's methylvalerate/95 mole's butyrate is introduced into a single screw extruder (Rheomix Model 202) with screw diameter of 0.75 cm.

ch. A constant larger screw having 20.1 length to demeter ratio and -31 compression ratio is employed. The temperature of both heating zones of the extruder barrel is 25°C above the met temperature of the PHBMV. The result of the screw of the screw

EXAMPLE 7 not example of the invention

Compostable Single Layer Film

[0149] Films of PHBMV (95:5) are made by melting the material between Teflon sheets in a Carver Press (Fred S. Carver Inc., Menomonee Falls, WI) at 20°C above the melt temperature. Pressure on the sheets are adjusted to produce films of approximately 0.25 mm thick. The films are then identically cooled to room temperature by placing the molds between large (5 kg) aluminum plates and allowing the films to cool to room temperature.

EXAMPLE 8 not example of the invention

35 Compostable Multilayer Film

[0150] Sheets of PHBMV film may be prepared as in Example 6 of compositions PHBMV (65.5) and PHBMV (55.50). These sheets may then encase a sheet of a polymer with good oxygen barrier properties but a poor valer vapor transmission rate, or a polymer film that may be water soluble such a polyvierny alcoholy (PVA). The films are placed in carver press stacked in the following order PHBMV(95.5), PHBMV(50.50), PVA, PHBMV(50.50), PHBMV(95.5). The material is then pressed at a temperature 5°C above the melt temperature of PHBMV(95.50), and it is libelow the melting temperature of the PHBMV(95.5). After compression at 2000 lb for 30 min, the pressure is released and the film is allowed to cool to room temperature.

45 FXAMPLE 9

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Compostable Disposable Diaper

[0151] A disposable baby diaper according to this invention is prepared as follows. The dimensions listed are for a diaper intended for use with a child in the 6-10 kilogram size range. These dimensions can be modified proportionately for different size children, or for adult incontinence briefs, according to standard practice.

- Backsheet: 0.020 0.038 mm film consisting of a 92.8 poly(3-hydroxybulyrate-co-3-hydroxy-4-methylvalerate) copolymer (prepared as described in Example 1); width at top and bottom 33 cm; notched inwardly on both sides to a width-at-center of 28.5 cm; length 50.2 cm.
- Topsheet: carded and thermally bonded staple-length polypropylene fibers (Hercules type 151 polypropylene);
 width at top and bottom 33 cm; notched inwardly on both sides to a width-at-center of 28.5 cm; length 50.2 cm.
- 3. Absorbent core: comprises 28.6 g of cellulose wood pulp and 4.9 g of absorbent gelling material particles (com-

mercial polyacrylate from Nippon Shokubai); 8.4 mm thick, calendered; width at top and bottom 28.6 cm; notched inwardly at both sides to a width-at-center of 10.2 cm; length 44.5 cm.

 Elastic leg bands: four individual rubber strips (2 per side); width 4,77 mm; length 370 mm; thickness 0.178 mm (all the foregoing dimensions being in the relaxed state).

[0152] The diaper is prepared in standard fashion by positioning the core material covered with the topsheet on the backsheet and cluing.

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FXAMPLE 10

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Compostable Lightweight Pantiliner

[0144] A lightweight pentiliner suitable for use between menstrual periods comprises a pad fourface area 117 cm². SSK air fet 3.0 g) containing 1.0 g of absorbert geilling material particles (commercial polyacystars, Nippon Shoukush) said pad being interposed between a porrus formed-fills topshet according to U.S. Patent 4.483,045 and a backsheet which comprises a 0.03 mm thickness 928 poly(3-hydroxybulyrate-co-3-hydroxy-4-methylvalerate) copolymer copolymer film, as prepared in accordinace with Example 3.

FXAMPLE 11

Compostable Sanitary Napkin

[0155] A catamenial product in the form of a sanitary napkin having two flaps extending outward from its absorbent core is prepared using a pad in the manner of Example 10 (surface area 117 cm²; 8.5 g SSK air felt), per the design of U.S. Patent 4,687.479, Van Tillburg. August 18, 1987. The backsheet and lopsheet materials are the same as described in Example 10.

EXAMPLE 12

Compostable Sheet

[0156] The film preparation procedure of Example 6 is modified by replacing the die on the extruder with a slot die of thickness approximately 0.25 cm and width 15 cm. Take-up after extrusion is accomplished by inserting the sheet emerging from the extruder between two counter-rotating cylinders. The sheet is drawn from the extruder in this manner and cut in lengths of 32 cm. Sheets of approximately 13 cm wide and 0.18 cm thick are obtained.

45 EXAMPLE 13

Compostable Fiber

[0157] PHBMV of composition 5 mole% methylvalerate/95 mole% butyrate is introduced into a single screw extruder (Rheomix Model 202) with screw diameter of 0.75 inch. A constant taper screw having 20.1 length to diameter ratio and 31 compression ratio is employed. The temperature of both heading zones of the extruder barrel is 25°C above the melt temperature of the PHBMV. The extruder is equipped with a nozzle die containing 5 ordices of diameter 500 mm. The die is maintained at 20°C above the melt temperature of the PHBMV. The polymer is melted within the extruder and pumped to the die at the other end of the extruder. The screw rpm is kept constant at 30 rpm. The polymer is forced through the de and the melted extruded fibers are lead through a region where a rapid ai stream is applied such that he polymer tibers elongates and thins to approximately one fith of the diameter of the ordices (ca. 100 mm). The fibers are collected on a carbidorad mail. A wide distribution of fiber lengths are obtained up several cm in length. Most fiber lengths (over 50%) are in the range of 1.3 to 15°C.

EXAMPLE 14

Compostable Rigid Foam

[0158] PHBMV (40 g) of composition 5 mole% methylvalerate/95 mole% butyrate and 4 g of a common blowing agent, p.p.-cvy-bis benzenesus/biorhydrazide are charged to the mixing chamber of a Rheomix type 600 meth bender euipped with roller blades. The mixing chamber temperature is heated above the mething temperature of PHBMV, but below the degradation temperature of the blowing agent (156°C). After mixing for 10 minutes at 60 pm, the copolymer mixture is collected and is transferred to a heated aluminum pan, spread about so that the resulting mass is about 0.5 cm in trickness. The copolymer is then place in an oven (National Appliance Company, model 5830) and heated to the PHBMV met temperature again, and is held at that temperature until the copolymer is completely molten (ca. 5 min). The oven temperature is then resised to 160°C at which temperature until the copolymer is completely molten (ca. 5 min). The oven Chamber is then resised to 160°C at which temperature the blowing agent degrades and copolymer begins foaming. At this point the copolymer foam is removed from the oven and is placed into a second over at a temperature of the maximum crystalization rate of the PHBMV (about 20°C). The copolymer is 6 hours.

EXAMPLE 15

Compostable Flexible Foam

[0159] The procedure of Example 14 is used with the following modifications: 40 g of poly(3-hydroxybulyrate-co-3-hydroxy4-methylvalerate) copolymer of composition 60 mole% methylvalerate/40 mole% butyrate (PHBMV (40:60)) is used in place of PHBMV (40:50).

EYAMPLE 16

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Compostable Molded Article

[0160] Injection molded articles are obtained by using a Mini Max Molder model CS-183 (Custom Scientific Instruments, Whippeny, N.J.). The temperature of the rotor and strator cup is held constant at 20°C above the melt temperature of the polyhydroxyalkanoate used. About 0.5 grams of PHBMV (95.5) is charged to the stator cup and allowed to melt for 3 minutes. The molten copolymer is radially mixed by raising and lowering the rotor tip five times. A dumbbell-shaped steel mold is sprayed with a light cealing of mold stillone release apent. The mold is placed on the mold support wheel of the Mini Max Molder and the molten polymer is injected into the mold by action of the rotor tip. The copplymer is molded into a dumbbell shaped pieces 0.03 inch thick, I inch long, 0.125 inch wide at the middle of the piece and 0.25 inch wide at the ends. These molded parts are suitable for mechanical testing.

EXAMPLE 17

Compostable Nonwoven Fabric

[0161] Poy(3-hydroxybutyrate-co-3-hydroxy-4-methylvalerate) copolymer (PHBMV) of composition Z mole's methylvaleratel' 98 mole's butyrate is introduced into a single screw acturder (Rheomix Model 202, Paramus, NJ) with screw
diameter of 0.75 inch. A constant taper screw having 20:1 length to diameter ratio and a 3:1 compression ratio is
employed. The temperature of both heating zones of the extruder barrel is 25°C above the melt temperature of the
PHBMV. The extruder is equipped with a nozzie die containing 5 onlines of diameter 500 mm. The die is mantalend at
20°C above the melt temperature of the PHBMV. The polymer is melted within the extruder rand pumped to the die at
the other end of the extruder. The screw pm is kept constant at 30 pm. The polymer is forced through the die and the
melted extruded fibers are lead through a region where a rapid air stream is applied such that the polymer fibers elongates
and thins to approximately one fifth of the diameter of the onlines (ac. 100 mm). The fibers are collected on a carboard
mat. The mat is moved in a fashion so that a 10 cm x 10 cm area is covered uniformly with fibers. Collection of fibers
on the mat continues, until there is approximately 0.5 cm thick fiber mat. A wide distribution of fiber lengths are obtained
up several inches in length. Most fiber lengths (sover 50%) are in the range of 0.5 to 6 inches. The mat is then transferred
to a Carver Press (Fred S. Carver Inc., Menomonee Falls, WI) and pressed at a 10001b force for 10 minutes at temperature
5°C below the melting temperature of the PHBMV. The resulting nonvoven sheet is removed from the press.

EXAMPLE 18

Compostable Elastomer

5 (0162) Films of PHBMV (70:30) are made by melting the material between Teflon sheets in a at 20°C above the melt temperature. Pressure on the sheets is adjusted to produce films of approximately 0.5 mm thick. The films are then identically cooled to room temperature by placing the molds between large (6.8) adminum plates and allowing the films to cool to room temperature. The films are sped for 2 days, then subsequently cut into strips 10 cm long and 1 cm wide. The strips are then placed in an instron universal testing materiale (Model 112/C, Canton, MA) and are elongated as 1 are led 1 invinin until 300% elongation of the original length is achieved. The films are held elongated for two days until crystallinity develops further. The strips are removed from the Instron and upon subsequent extension, the material returns to its formor (post Instron realment) length.

FXAMPLE 19

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Compostable Adhesive

[0163] PHBMV (50:50) may be used as a hot-melt adhesive in the following manner. About 1g of PHBMV (50:50) is placed believen two polymer films, such as poly(vinyl alcohol) (PVA), or poly(3-hydroxybutyste) (PHB) or any other PHA which has a melting temperature at least 10°C higher than PHBMV (50:50). The films/adhesive assembly is placed in a Carver Press (Fred S. Carver Inc., Menomonee Falls, WI) and is then pressed at a temperature 5°C above the melt temperature of PHB.MV (50:50). After compression at 2000 to force for 30 min, the pressure is released and the bonded film assembly is allowed to cool to room temperature.

101641 All publications mentioned hereinabove are hereby incorporated in their entirety by reference.

[0165] It is understood that the examples and embodiments described herein are for illustrative purposes only and that various modifications or changes in light thereof will be suggested to one skilled in the art and are to be included in the spirit and purview of this application and scope of the appended claims.

30 Claims

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 A plastic article other than a film comprising a biodegradable copolymer, wherein the biodegradable copolymer comprises at least two randomly repeating monomer units wherein the first randomly repeating monomer unit has the structure.

wherein R1 is H, or C1 or C2 alkyl, and n is 1 or 2; the second randomly repeating monomer unit has the structure

- and wherein at least 50% of the randomly repeating monomer units have the structure of the first randomly repeating monomer unit.
- The plastic article of Claim 1, wherein R¹ is a C₁ or C₂ alkyl and n is 1.

- The plastic article of Claim 2, wherein R¹ is a C₁ alkyl.
- The plastic article of Claim 1, wherein R¹ is H and n is 2.
- The plastic article of Claim 1, wherein the plastic article is a fiber.
 - 6. The plastic article of Claim 1, wherein the plastic article is a foam.
 - 7. The plastic article of Claim 1, wherein the plastic article is a molded article.
 - 8. The plastic article of Claim 1, wherein the plastic article is a nonwoven fabric.
- The plastic article of Claim 1, wherein the plastic article is an elastomer.
- 15 10. The plastic article of Claim 1, wherein the plastic article is an adhesive.
 - The plastic article of Claim 1, wherein the plastic article is a sheet.
 - The plastic article of Claim 1, wherein the copolymer comprises one or more additional randomly repeating monomer units having the structure

- 30 wherein R³ is H, or a C₁-C₁₉ alkyl or alkenyl; and m is 1 or 2; and wherein the additional randomly repeating monomer units are not the same as the first randomly repeating monomer unit or the second randomly repeating monomer unit.
- 13. The plastic article of Claim 12, wherein the plastic article is a fiber.
- 5 14. The plastic article of Claim 12, wherein the plastic article is a foam.
 - 15. The plastic article of Claim 12, wherein the plastic article is a molded article.
 - The plastic article of Claim 12, wherein the plastic article is a nonwoven fabric.
 - 17. The plastic article of Claim 12, wherein the plastic article is an elastomer.
 - 18. The plastic article of Claim 12, wherein the plastic article is an adhesive.
- 45 19. The plastic article of Claim 12, wherein the plastic article is a sheet.

Revendications

 Article en plastique autre qu'une pellicule comprenant un copolymère biodégradable, dans lequel le copolymère biodégradable comprend au moins deux moilts monomères rojetés de manière aléatoire dans lesquels le premier moif imnommère répété de manière aléatoire a la structure

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dans laquelle \mathbb{R}^1 est H, ou un alkyle en \mathbb{C}_1 ou \mathbb{C}_2 , et n est 1 ou 2; le deuxième motif monomère se répétant de manière aléatoire a la structure

et dans laquelle au moins 50 % des motifs monomères se répétant de manière aléatoire ont la structure du premier motif monomère se répétant de manière aléatoire.

- Article en plastique de la revendication 1, dans lequel R¹ est un alkyle en C₁ ou C₂ et n est 1.
 - Article en plastique de la revendication 2, dans lequel R¹ est un alkyle en C₁.
 - Article en plastique de la revendication 1, dans lequel R¹ est H et n est 2.

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- Article en plastique de la revendication 1, dans lequel l'article en plastique est une fibre.
 - 6. Article en plastique de la revendication 1, dans lequel l'article en plastique est une mousse.
 - Article en plastique de la revendication 1, dans lequel l'article en plastique est un article moulé.
 - 8. Article en plastique de la revendication 1, dans lequel l'article en plastique est un tissu nontissé.
 - Article en plastique de la revendication 1, dans lequel l'article en plastique est un élastomère.
 - Article en plastique de la revendication 1, dans lequel l'article en plastique est un adhésif.
 - 11. Article en plastique de la revendication 1, dans lequel l'article en plastique est une feuille.
- 45 12. Article en plastique de la revendication 1, dans lequel le copolymère comprend un ou plusieurs motifs monomères se répétant de manière aiéatoire supplémentaires ayant la structure

dans laquelle R³ est H, ou un alkyle ou alcényle en C₁ à C₁₉; et m est 1 ou 2; et dans laquelle les motifs monomères se répétant de manière aléatoire supplémentaires ne sont pas les mêmes que le premier motif monomère se répétant de manière aléatoire ou le deuxième motif monomère se répétant de manière aléatoire.

- Article en plastique de la revendication 12, dans lequel l'article en plastique est une fibre.
- 14. Article en plastique de la revendication 12, dans lequel l'article en plastique est une mousse.
 - 15. Article en plastique de la revendication 12, dans lequel l'article en plastique est un article moulé.
 - Article en plastique de la revendication 12, dans lequel l'article en plastique est un tissu nontissé.
 - 17. Article en plastique de la revendication 12, dans lequel l'article en plastique est un élastomère.
 - Article en plastique de la revendication 12, dans lequel l'article en plastique est un adhésif.
- 19. Article en plastique de la revendication 12, dans lequel l'article en plastique est une feuille.

Patentansprüche

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1. Anderer Kunststoffartikel als ein Film, der ein biologisch abbaubares Copolymer umfasst, worin das biologisch abbaubare Copolymer mindestens zwei statistisch wiederkehrende Grundmonomereinheiten umfasst, worin die erste statistisch wiederkehrende Grundmonomereinheit die Struktur

aufweist, worin R1 H oder C1- oder C2-Alkyl ist und n 1 oder 2 ist; die zweite statistisch wiederkehrende Grundmonomereinheit die Struktur

- aufweist und worin mindestens 50 % der statistisch wiederkehrenden Grundmonomereinheiten die Struktur der ersten statistisch wiederkehrenden Grundmonomereinheit aufweisen.
- Kunststoffartikel nach Anspruch 1, worin R¹ ein C₁- oder C₂-Alkyl ist und n 1 ist.
- 3. Kunststoffartikel nach Anspruch 2, worin R1 ein C1-Alkyl ist.
 - 4. Kunststoffartikel nach Anspruch 1, worin R1 H ist und n 2 ist.
 - 5. Kunststoffartikel nach Anspruch 1, worin der Kunststoffartikel eine Faser ist.
- 6. Kunststoffartikel nach Anspruch 1, worin der Kunststoffartikel ein Schaumstoff ist.
 - Kunststoffartikel nach Anspruch 1, worin der Kunststoffartikel ein Formartikel ist.

- 8. Kunststoffartikel nach Anspruch 1, worin der Kunststoffartikel ein Vliesstoff i st.
- Kunststoffartikel nach Anspruch 1, worin der Kunststoffartikel ein Elastomer ist.
- Kunststoffartikel nach Anspruch 1, worin der Kunststoffartikel ein Haftmittel ist.
 - Kunststoffartikel nach Anspruch 1, worin der Kunststoffartikel eine Folie ist.

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 Kunststoffartikel nach Anspruch 1, worin das Copolymer eine oder mehrere zusätzliche statistisch wiederkehrende Grundmonomereinheiten mit der Struktur

umfasst, worin R³ H oder ein C₁-C₁₉-Alkyl oder-Alkenyl ist und m 1 oder 2 ist; und worin die zusätzlichen stallsitsch wiederkohrenden Grundmonomereinheiten richt dieselben sind wie die erste stalistisch wiederkehrende Grundmonomereinheit oder die zweite stalistisch wiederkehrende Grundmonomereinheit.

- Kunststoffartikel nach Anspruch 12, worin der Kunststoffartikel eine Faser ist.
- 14. Kunststoffartikel nach Anspruch 12, worin der Kunststoffartikel ein Schaumstoff ist.
- Kunststoffartikel nach Anspruch 12, worin der Kunststoffartikel ein Formartikel ist.
- 30 16. Kunststoffartikel nach Anspruch 12, worin der Kunststoffartikel ein Vliesstoff ist.
 - 17. Kunststoffartikel nach Anspruch 12, worin der Kunststoffartikel ein Elastomer ist.
 - 18. Kunststoffartikel nach Anspruch 12, worin der Kunststoffartikel ein Haftmittel ist.
 - Kunststoffartikel nach Anspruch 12, worin der Kunststoffartikel eine Folie ist.